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# Single-atom alloy structure and unique bonding properties of Au<sub>104</sub>Ag<sub>40</sub>(PET)<sub>60</sub> nanoclusters†

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The detailed characterization of AuAg alloy nanoclusters is essential to guide the discovery of species ideal for applications in various fields including catalysis and biomedicine. This work presents structural analysis of the Au<sub>104</sub>Ag<sub>40</sub>(PET)<sub>60</sub> species through X-ray absorption spectroscopy (XAS). First, XAS fitting is utilized to model the distribution of Au and Ag atoms within the structure. Our proposed model assigns Ag atoms to the vertex sites of the second shell of the metal core, as well as the outermost staple sites. This distribution reveals Au<sub>104</sub>Ag<sub>40</sub>(PET)<sub>60</sub> to be a Ag single-atom alloy. The proposed model shows outstanding agreement with the coordination number values derived from XAS. XAS near-edge analysis is employed to investigate the alloy bonding interactions between Au and Ag. Substantial d-electron transfer from Au to Ag is observed in this sample, beyond the magnitude of previously studied AuAg NCs. This work enhances the understanding of the structure–property relationship of AuAg alloy NCs, offering insights which can be applied to other large NCs and even NPs. These insights will in turn aid the discovery of new materials for use in various applications.

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#### 1. Introduction

Au nanoclusters (NCs) have been widely studied due to the many unique properties which arise from their ultrasmall size. These properties result in promising candidates for a wide variety of applications, in fields including catalysis, optics, and electronics. As the properties of a NC are dependent on structural features including shape, size, and atomic arrangement, detailed characterization is vital to establish a structure–property relationship and help guide the discovery of NC species with practical applications. To this end, atomically precise NCs, defined by their uniform, well-defined structure, have emerged as an area of focus due to their consistent and predictable properties. 5-9

Beyond monometallic NCs, introducing a second metal to form alloy NCs increases the compositional space, unlocking many new possibilities. <sup>10–12</sup> By using atomically precise NCs as a starting template the properties can be carefully tuned through adjusting the exact dopant concentration of the second metal, which in turn modifies the composition of the final product. This high degree of control increases the feasibility of creating alloy NCs to be utilized for desirable applications. <sup>13</sup> This includes classes of materials such as single-

atom alloys, which are of interest due to their excellent catalytic activity and selectivity. 14-19

One widely studied combination of metals is Au and Ag, owing to the synergistic effects between them resulting in enhanced catalytic performance, stability, biocompatibility. 10,11,20 AuAg alloy NCs have been successfully synthesized through methods including galvanic replacement and co-reduction.<sup>20</sup> The exact atomic positions that each element occupies has been shown to have a significant impact on the properties expressed by the NC.21 Therefore, an accurate determination of these positions is vital to develop the structure-property relationship. Previous work has accomplished this for relatively large AuAg NCs including (AuAg)<sub>130</sub> and (AuAg)<sub>267</sub>.<sup>21,22</sup> These studies primarily relied on X-ray crystallography, which requires the synthesis of a high-quality single-crystal. This has proven to be very challenging due to the complex and flexible structures of NCs.<sup>23</sup> X-ray crystallography data interpretation can be challenging for large NCs due to the high atomic number contrast between heavy metal atoms in the core and lighter atoms in the thiolate ligands, leading to uncertainty in the atomic identity and exact atomic positions. 21,24 Other techniques also run into challenges for the analysis of large AuAg NCs. Imaging techniques struggle to observe the metal core as it is covered by the protecting organothiolate ligands, and while DFT calculations can provide useful information, the resources necessary to run these simulations increase significantly when considering NCs of increased size and multiple elements.5

A tool that has shown great success in the characterization of metal NCs is X-ray absorption spectroscopy (XAS). 4,8,10,11,25,26 XAS

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allows for flexible sample preparation, circumventing the single-crystal requirement for X-ray crystallography. XAS provides element specific analysis, making it ideal for alloy samples containing multiple elements. X-ray absorption near-edge spectroscopy (XANES) provides insight into electronic properties, and extended X-ray absorption fine structure (EXAFS) yields detailed structural information including accurate values for the average coordination number (CN) and bond distances (*R*) of various interactions within the sample.<sup>27–29</sup> All of these features makes XAS an ideal tool to provide insight into the structure–property relationship of metal alloy NCs.

This work focuses on the previously reported<sup>30</sup> Au<sub>104</sub>Ag<sub>40</sub> NC, utilizing a XAS-focused characterization strategy to gain an in-depth understanding of the structure of this sample. The previously reported crystal structure of Au<sub>144</sub>,<sup>24</sup> the monometallic analogue of Au<sub>104</sub>Ag<sub>40</sub>, is used as a template to assign likely Ag locations, resulting in a model representing the exact structure of Au<sub>104</sub>Ag<sub>40</sub>. Our proposed model assigns 40 Ag atoms within the 144-metal atom structure, with 30 of them occupying the entirety of the staple sites and the remaining 10 occupying vertex sites of the second shell of the metal core. This model shows excellent agreement with the EXAFS fitting results, as well as consistent trends with other AuAg NCs of similar size.21,22 Excitingly, our findings indicate that Au<sub>104</sub>Ag<sub>40</sub> is a single-atom Ag alloy, making it a promising candidate for applications such as catalysis. XANES analysis is utilized to probe the electronic properties of the sample, indicating d-electron transfer away from Au at a larger magnitude than what has been observed in other NCs of similar size.

## 2. Experimental methods

#### 2.1 Reagents and materials

Tetrachloro Auric(III) acid (HAuCl<sub>4</sub>·3H<sub>2</sub>O, >99.99% metals basis, Aldrich), silver tetrafluoroborate (AgBF<sub>4</sub>, >99.99% metals basis, Aldrich), tetraoctylammonium bromide (TOAB,  $\geq$ 98%, Fluka), 2-phenylethanethiol (PET, C<sub>8</sub>H<sub>9</sub>SH, 98%, Aldrich), sodium borohydride (NaBH<sub>4</sub>, Aldrich), methanol (MeOH, HPLC grade,  $\geq$ 99.9%, Aldrich), dichloromethane (DCM, ACS reagent,  $\geq$ 99.5%, Aldrich). All chemicals were used without further purification. Nanopure water was prepared using the Barnstead NANOpure Diamond system.

#### 2.2 Synthesis and mass spectrometry

Au<sub>104</sub>Ag<sub>40</sub> NCs were synthesized following previously reported procedures.<sup>30</sup> Briefly, 120 mg of HAuCl<sub>4</sub>·3H<sub>2</sub>O and 30 mg of AgBF<sub>4</sub> were dissolved in 10 mL of methanol and vigorously stirred for 15 minutes. Under continued vigorous stirring, 190 mg of 2-phenylethanethiol in the presence of tetraoctylammonium bromide was added to the reaction mixture, followed by 114 mg of NaBH<sub>4</sub> dissolved in 5 mL of cold Nanopure water. Black precipitate containing the NCs was precipitated out of solution. The remaining solution was stirred overnight, after which the supernatant was separated *via* centrifuge and discarded. The remaining precipitate was washed three times

with methanol, then NCs were extracted using dichloromethane. The exact mass of the product was determined through electrospray ionization mass spectrometry (ESI-MS), collected using a Waters Q-TOF mass spectrometer equipped with Z-spray source.

#### 2.3 X-ray absorption spectroscopy

Au L3-edge and Ag K-edge XAS data was collected from the CLS@APS (Sector 20-BM) beamline at the Advanced Photon Source (operating at 7.0 GeV) in Argonne National Laboratory, Chicago, IL, USA. Solid powdered sample was packed into kapton tape and measured in fluorescence mode simultaneously with a metal foil reference for each element. All measurements were collected at room temperature under ambient pressure. EXAFS data was transformed and normalized into k-space (Fig. S1†) and R-space using the Athena program following conventional procedures.31 A k-range of  $2.9-13.4 \text{ Å}^{-1}$  for Au, and  $3.4-10.5 \text{ Å}^{-1}$  for Ag was used to obtain each Fourier-transformed EXAFS (FT-EXAFS) spectra with a k weighting of 2. Self-consistent multiple-scattering calculations were performed using the FEFF6 program to obtain the scattering amplitudes and phase-shift functions used to fit scattering paths within the Artemis program. In the fitting a R-window of 1.2-3.6 Å for Au and 1.5-3.6 Å for Ag was used. Au L<sub>3</sub>-edge and Ag K-edge data were simultaneously fitted, and various parameters were correlated together to minimize the number of independent variables and ensure a high fit quality. These correlations are clearly indicated in the relevant data tables.

### 3. Results and discussion

#### 3.1 Structural determination of Au<sub>104</sub>Ag<sub>40</sub>

Au<sub>104</sub>Ag<sub>40</sub> NCs were synthesized following the previously reported procedure. The sample of interest in this work was synthesized using a 2:1 molar ratio of Au<sup>III</sup>/Ag<sup>I</sup>. ESI-MS results (Fig. S2†) determine the most abundant species within this sample contains 40 Ag and 104 Au atoms. The well-defined peaks in the ESI-MS spectrum provide confidence in the purity of Au<sub>104</sub>Ag<sub>40</sub>. As such, we assume the identity of Au<sub>104</sub>Ag<sub>40</sub> to be accurate, and that the shape and atomic positions of Au<sub>104</sub>Ag<sub>40</sub> will be consistent with the Au<sub>144</sub> structure determined through X-ray crystallography. Our approach is to use the structural parameters calculated from EXAFS fitting to determine the most likely locations that the 40 dopant Ag atoms will occupy within this 144-metal atom structure.

The metal atoms in Au<sub>144</sub> can be separated into 4 categories, each of which forms a "shell" within the structure. Beginning at the innermost core of the NC, shell 1 contains 12 atoms arranged in icosahedral geometry, with the notable lack of a center atom resulting in a M-M CN of 11 for each site. Shell 2 contains 42 atoms, also in icosahedral geometry. The 12 vertices of shell 2 have a M-M CN of 11, while the remaining shell 2 atoms have a M-M CN of 12. Shell 3, which forms the surface of the NC, has a rhombicosidodecahedron shape and contains 60 atoms, each with a M-M CN of 9. Finally shell

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4, which takes the form of a series of staple units, contains 30 atoms, each with a M-M CN of 4. The S atoms in the protective organothiolate ligands form bonds with M atoms in shells 3 and 4. Atoms in each shell will form M-M bonds with atoms in the shell below and above it, as well as bonds with other atoms within the same shell. The exception is shell 4, which will only form bonds with shell 3 atoms due to the long distance between shell 4 atoms. The Au<sub>144</sub> bond distribution (Fig. S3†) shows a general trend of shorter M-M bonds with an approximate bond length ~2.8 Å towards the core, and longer M-M bonds with an approximate bond length ~3.0 Å towards the surface. With this understanding of the structure of Au<sub>144</sub> established, EXAFS fitting was conducted on the Au<sub>104</sub>Ag<sub>40</sub> to compare structural parameters to those observed in Au<sub>144</sub>.

Wavelet transform-EXAFS (WT-EXAFS) plots (Fig. 1) were used to observe the most intense features within the k-space and R-space, allowing for an appropriate fitting procedure to be chosen for the EXAFS fitting procedure. Intense features in the WT-EXAFS plot indicate distinct bonding interactions which should have separate fitting paths in the EXAFS fitting procedure.<sup>32</sup> Feature A represents a Au-S interaction, and feature D represents a Ag-S interaction. As both Au and Ag show evidence of a significant bonding interaction with S atoms, it is concluded that both Au and Ag must occupy positions within shell 3 and/or shell 4. Two additional distinct maxima are observed in both the Au L3-edge (features B and C) and the Ag K-edge (features E and F). These features are within the region characteristic of M-M interactions. The presence of two distinct features indicates that there is a wide distribution of the Au-M and Ag-M bond lengths, consistent with what is observed in Au<sub>144</sub> (Fig. S3†). As both Au and Ag show evidence of a wide range of M-M bond distances, it is likely that they are both distributed relatively evenly across the structure, ruling out the possibility of a core-shell type structure.

Based on the features observed in the WT-EXAFS plots, a multi-shell fitting method for the M-M bonds is necessary to accurately model this system. The fitted FT-EXAFS spectra (Fig. S4†) and EXAFS fitting results (Table 1) allow for the average CN values of various bond types to be calculated. The most reliable fitting result utilizes four fitting paths for Au (Au-S, Au-Au, 2× Au-Ag) and three fitting paths for Ag (Ag-S, 2× Ag-Au). Interestingly, all fitting attempts which used a Ag-Ag fitting path gave a poor fit quality and infeasible values such as a negative CN. This result is strong evidence of a lack of Ag-Ag nearest-neighbor interactions in the sample, making Au<sub>104</sub>Ag<sub>40</sub> a single-atom Ag alloy.

Given the large proportion of Ag in this sample, there are very few possible configurations which will satisfy a Ag singleatom alloy distribution. The lack of nearest-neighbor Ag atoms is the main guiding principle for the proposed model, but there are also many other conditions which must be satisfied. The Au-Au fitting path, with a relatively large CN of 6 and relatively short bond distance of 2.81 Å, indicates a Au-rich metal core. The two distinct Au-Ag fitting paths indicate Au-Ag interactions are present within shells 1 and 2 (2.84 Å) and within shells 3

Table 1 EXAFS fitting results for the  $Au_{104}Ag_{40}$  sample. Data for the Au $L_3$ -edge and Ag K-edge was simultaneously fit. R values for the Au-Aq and Aq-Au fitting paths were correlated as they represent the same chemical bond, and therefore must have the same bond distance. Debye-Waller ( $\sigma^2$ ) values for the Au-M fitting paths, and for all the Ag fitting paths were correlated to minimize the number of independent variables and ensure a high-quality fit.  $E_0$  values for the Au fitting paths and Ag fitting paths were also correlated

Bond	CN	R (Å)	$\sigma^2 \left(\mathring{A}\right) \times 10^{-3}$	$E_0$ (eV)	R factor
Au-S Au-Au Au-Ag Au-Ag	0.9(2) 6(2) 1.0(9) 1.4(8)	2.30(1) 2.81(2) 2.84(4) 2.99(4)	5(2) 17(4)	7(1)	0.0249
Ag-Ag Ag-Au Ag-Au	0.8(4) 3(1) 3(1)	2.43(3) 2.84(4) 2.99(4)	4(5)	1(3)	0.0249

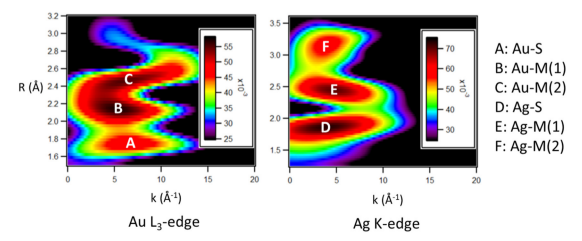


Fig. 1 WT-EXAFS plots for  $Au_{104}Ag_{40}$ . Intense features are labelled according to the bond types they represent. In the  $Au\ L_3$ -edge spectra: feature  $Au\ L_3$ -edge spectra: feature represents Au-S, feature B represents the shorter distance Au-M, and feature C represents the longer distance Au-M. In the Aq K-edge: feature D represents Ag-S, feature E represents the shorter distance Ag-M, and feature F represents the longer distance Ag-M.

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and 4 (2.99 Å). The Au-S and Ag-S CNs are roughly equal, providing insight into the elemental distribution of shells 3 and 4. Finally, the total Ag-M CN (6) is lower than that of Au (8.4). This finding further supports the innermost part of the structure being dominated by Au. It also indicates that shell 4, having a low M-M CN of 4, will be largely occupied by Ag.

Accounting for all these structural features, a model has been proposed for the exact element distribution of Au<sub>104</sub>Ag<sub>40</sub> (Fig. 2). The CN values of this model show remarkable agreement with those calculated from EXAFS (Table 2), particularly for the alloy bonding interactions, providing high confidence in this assignment. Note that it is understandable to have a high degree of uncertainty for the M-S interactions due to the large NC size. In this model, all 30 of the shell 4 sites are occupied with Ag, and all 60 of the shell 3 sites are occupied with Au. This assignment satisfies a single-atom Ag distribution, a low overall Ag-M CN, and a roughly equal CN of Au-S and Ag-S. The staple layer being dominated by Ag has precedent in other AuAg NCs. 21 Furthermore, the truncated geometry of the shell 3 surface layer has been observed in many monometallic Au NCs, as a way to reduce the high surface energy of Au, while monometallic Ag NCs have shown preference for a nontruncated geometry.33,34 Therefore, in the case of a surface shell with truncated shape as in Au<sub>104</sub>Ag<sub>40</sub>, Au showing a preference for occupying these shell 3 sites is expected.

The remaining 10 Ag atoms have been assigned to the vertex sites of the icosahedron of shell 2. While other assignments of Ag within shell 2 satisfy the CN and bond distance values determined through EXAFS, this specific assignment is justified based on a similar distribution observed in the (AuAg)<sub>130</sub> NC through X-ray crystallography.<sup>21</sup> The top and bottom vertices of the shell 2 icosahedron were chosen to be the two vertices that are occupied by Au in our model. This is necessary to ensure a model containing exactly 40 Ag atoms, but also has precedent in  $(AuAg)_{130}$ .<sup>21</sup>

To validate this model, the EXAFS fitting was refined with the CN values fixed according to the values in the model (Fig. S5<sup>†</sup> and Table 3). A high-quality fit is obtained using this fitting method, providing confidence in the proposed model. By fixing the CN values more independent variables can be utilized in the fitting procedure, allowing for the uncorrelation of

Table 2 CN comparison between experimental EXAFS fitting results and the proposed model for the  $Au_{104}Ag_{40}$  sample

Bond	CN (EXAFS)	CN (model)	
Au-S	0.9(2)	0.6	
Au-Au	6(2)	7.7	
Au-Ag	1.0(9)	1.0	
Au-Ag	1.4(8)	1.2	
Ag-S	0.8(4)	1.5	
Ag-Au	3(1)	2.8	
Ag-Au	3(1)	3.0	

Table 3 Refined EXAFS fitting results for the Au<sub>104</sub>Ag<sub>40</sub> sample. CN values are fixed according to values from the proposed model. Data for the Au L<sub>3</sub>-edge and Ag K-edge was simultaneously fit. R values for the Au-Ag and Ag-Au fitting paths were correlated as they represent the same chemical bond, and therefore must have the same bond distance.  $\sigma^2$  values for the Au-Au and first Au-Ag fitting paths were correlated to minimize the number of independent variables and ensure a highquality fit.  $E_0$  values for the Au fitting paths and Ag fitting paths were also correlated

Bond	CN (fixed)	R (Å)	$\sigma^2  (\mathring{A}) \times 10^{-3}$	$E_0$ (eV)	R factor
Au-S	0.6	2.28(1)	2(1)	5(1)	0.0209
Au-Au	7.7	2.81(2)	18(2)		
Au-Ag	1.0	2.85(2)	,		
Au-Ag	1.2	2.98(2)	8(1)		
Ag-S	1.4	2.45(2)	10(3)	2(1)	0.0209
Ag-Au	2.8	2.85(2)	5(3)		
Ag-Au	3.0	2.98(2)	6(3)		
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 $\sigma^2$  values which was previously necessary to ensure a highquality fit. This fitting method allows for further insight regarding the bond distance distribution of each fitting path in Au<sub>104</sub>Ag<sub>40</sub>. Notably the Au-Au and short-length Au-Ag bonds have a very large  $\sigma^2$  of  $18 \times 10^{-3}$  Å, indicating a wide distribution of these bond types in Au<sub>104</sub>Ag<sub>40</sub>.

#### 3.2 Alloy bonding of Au<sub>104</sub>Ag<sub>40</sub>

Following the discovery of the single-atom Ag nature of Au<sub>104</sub>Ag<sub>40</sub>, we conducted an in-depth investigation into the alloy bonding interactions present within this sample. The 30 Ag atoms located within shell 4 are non-metallic in nature,

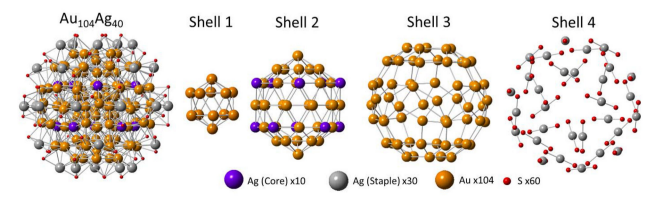


Fig. 2 Proposed model of Au<sub>104</sub>Ag<sub>40</sub>. Each shell is individually shown for clarity purposes.

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making them of less interest in applications such as catalysis. We therefore focus our analysis on the 10 Ag atoms localized within shell 2.

An in-depth analysis of this structure reveals several interesting features. Each of the Ag atoms has a CN of 11, with one shell 1, five shell 2, and five shell 3 neighbor atoms (Fig. 3). The EXAFS fitting results indicate Au-Ag bonds are on average slightly longer in length than Au-Au bonds. The lack of an innermost center atom in shell 1 provides the metal core of the NC with a higher degree of flexibility, allowing for the core to expand or contract as necessary to accommodate changes within the structure such as the introduction of dopant Ag atoms.

When considering Au<sub>104</sub>Ag<sub>40</sub> as a whole, this model can be viewed as having a "sandwich" type distribution of Au atoms, with shells 1 and 3 consisting entirely of Au and surrounding the shell 2 layer where the single-atom Ag sites are located. Au<sub>104</sub>Ag<sub>40</sub> approaches the upper limit of the number of Ag atoms which can be accommodated while still maintaining a single-atom alloy distribution. A high number of single-atom sites is of particular interest for applications such as catalysis, and this principle can potentially be applied to not only other AuAg NCs but also larger nanoparticles. Interestingly the final composition of this sample contained slightly less Ag than what was expected from the 2:1 Au: Ag loading, which may

indicate that additional dopant Ag atoms were repelled from the structure in order to ensure the single-atom Ag arrangement is maintained. This discrepancy indicates that Au NCs are a good template structure for the creation of single-atom Ag alloys.

Direct comparison with the similar sized (AuAg)<sub>130</sub> (Fig. 4) reveals interesting similarities and differences. As (AuAg)<sub>130</sub> has a roughly 1:2 ratio of Au: Ag, a single-atom Ag distribution is not possible. Despite this, the second shell shows a very similar distribution of Ag atoms to that of Au<sub>104</sub>Ag<sub>40</sub>, with Ag maintaining a single-atom alloy type distribution within shell 2. A similar phenomenon was also observed in the larger (AuAg)<sub>267</sub>, where the third shell (outermost core shell) also showed evidence of a single-atom alloy type Ag distribution.<sup>22</sup> The similarities in the distribution of Ag atoms within shell 2 hold despite the differences in shape of this layer between  $(AuAg)_{130}$  and  $Au_{104}Ag_{40}$ . When viewing  $(AuAg)_{130}$  from the side (Fig. 4a and c) the second shell can be divided into 7 horizontal rows of atoms, with the atoms in each row being arranged in the shape of a pentagon as can be clearly observed in the top-down view (Fig. 4b and d). The pentagon of every row has the same orientation, with all the vertices aligned.

For Au<sub>104</sub>Ag<sub>40</sub> the side view (Fig. 4e and g) also reveals 7 horizontal rows of atoms. While most of these rows maintain

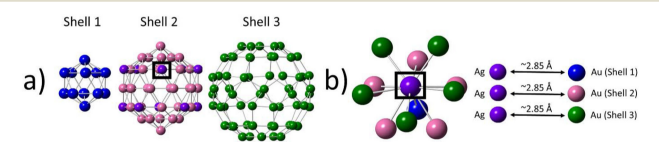


Fig. 3 Local bonding environment of the single-atom Ag sites within shell 2. (a) Breakdown of shells 1, 2, and 3. (b) Nearest-neighbor coordination environment of a single-atom Ag site in shell 2.

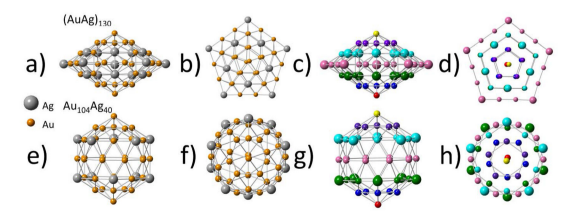


Fig. 4 Shell 2 comparison between (AuAg)<sub>130</sub> and Au<sub>104</sub>Ag<sub>40</sub>. (a) (AuAg)<sub>130</sub> side view. (b) (AuAg)<sub>130</sub> top-down view. (c) (AuAg)<sub>130</sub> side view (colored). (d)  $(AuAg)_{130}$  top-down view (colored). (e)  $Au_{104}Ag_{40}$  side view. (f)  $Au_{104}Ag_{40}$  top-down view. (g)  $Au_{104}Ag_{40}$  side view (colored). (h)  $Au_{104}Ag_{40}$  top-down view. down view (colored). Ag atoms are enlarged for clarity purposes. Colored view separates atoms by row to help highlight geometry from various viewpoints. In the top-down (colored) views, bonds between rows are omitted for clarity purposes.

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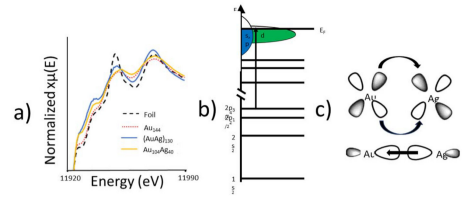


Fig. 5 Electronic properties of Au<sub>104</sub>Ag<sub>40</sub>. (a) Au L<sub>3</sub>-edge XANES spectra for Au<sub>104</sub>Ag<sub>40</sub> alongside metallic foil reference, Au<sub>144</sub>, and (AuAg)<sub>130</sub>. (b) Energy diagram illustrating the Au L<sub>3</sub>-edge electronic transition. (c) Schematic showing the electron transfer involved in the alloy interaction between Au and Ag atoms. Au donates d-electron density to Ag and receives s/p-electron density from Ag.

the same pentagon shape observed in (AuAg)<sub>130</sub>, the middle row instead has a decagon shape. From a top-down view (Fig. 4f and h) it is observed that rows in the top half have reverse orientation to rows in the bottom half. This mismatch results in shell 2 having a more rounded shape in Au<sub>104</sub>Ag<sub>40</sub>, and a more flat, spread-out shape in (AuAg)<sub>130</sub>. These changes in geometry are not enough to impact the bond distances, as EXAFS fitting indicates both (AuAg)<sub>130</sub> and Au<sub>104</sub>Ag<sub>40</sub> to have Au-Ag bond distances within uncertainty.

Interestingly, despite the 1:2 ratio of Au: Ag in (AuAg)<sub>130</sub>, this sample does not approach a single-atom Au type structure, with Au clustering in shell 2 and Ag spreading out across all 4 shells. This variation in the distribution between Au and Ag makes Ag the more likely candidate to form single-atom alloys at the appropriate composition. Comparison between (AuAg)<sub>130</sub> and Au<sub>104</sub>Ag<sub>40</sub> indicates that as the composition is varied to have a decreased amount of Ag, there will be less Ag atoms present within the innermost core (shell 1) and on the surface (shell 3) of the NC. Future study is necessary to confirm if this trend can be applied more broadly to AuAg NCs.

XANES was utilized to probe the electronic properties of the alloy bonding interaction. The Au L<sub>3</sub>-edge XANES spectra (Fig. 5a) of Au<sub>104</sub>Ag<sub>40</sub> shows an increased white-line intensity compared to the metallic foil reference. There are two plausible explanations for the increase white-line intensity in Au<sub>104</sub>Ag<sub>40</sub>; the nanoscale effect, or a decrease in d-electron density given that the L<sub>3</sub>-edge represents an electronic transition to the valence d-orbital (Fig. 5b). As Au<sub>144</sub> shows a similar white-line intensity to the metallic foil, the nanoscale effect can be ruled out and it can be concluded that a decrease in d-electron density is responsible. This finding indicates that Au donates d-electrons to Ag to form the alloying interaction, and in turn receives s/p-electron density back from Ag (Fig. 5c). Similar charge transfer has been previously observed in AuAg<sub>130</sub>.<sup>21</sup>

To quantitatively evaluate the alloying phenomenon, XANES integration was used to calculate the d-hole (10 -

Table 4 Calculated Au d-hole from XANES integration

Sample	d-Hole
Au <sub>144</sub>	0.372
Au foil	0.400
(AuAg) <sub>130</sub>	0.413
Au <sub>104</sub> Ag <sub>40</sub>	0.673

number of d electrons) of Au for each of these samples (Table 4). The d-hole for Au<sub>144</sub> is slightly smaller than for Au foil, as expected due to quantum size effects. Despite being of similar size, both  $(AuAg)_{130}$  and  $Au_{104}Ag_{40}$  have a larger d-hole than the Au foil, arising from d-electron density being transferred to Ag from Au. Interestingly, the single-atom Ag structure of Au<sub>104</sub>Ag<sub>40</sub> results in a substantially larger d-hole, due to Au-S interactions which are not present in (AuAg)<sub>130</sub>. The relative charge of the catalytically active site has been shown to play a key role in the catalytic performance of alloy NCs, making this observation of particular importance for future catalytic applications. 35,36

#### 4. Conclusions

XAS analysis has been used to identify the exact locations occupied by Au and Ag within the  $Au_{104}Ag_{40}$  NC. In this model Ag occupies all 30 staples sites and 10 of the 12 vertices of the second shell of the metal core, resulting in a Ag single-atom alloy structure. The multi-step EXAFS fitting procedure provides a high degree of confidence for this model, showcasing how structural insight can be obtained for large AuAg NCs when single-crystal synthesis is not possible. Structural analysis revealed Au<sub>104</sub>Ag<sub>40</sub> is an example of a single-atom Ag alloy, with Ag atoms located in a "sandwich" distribution between layers of Au atoms. XANES analysis provided insight into the alloy interaction between Au and Ag, revealing that Au donates d-electron density to Ag and receives s/p-electron

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density from Ag. This understanding of the charge transfer between elements is of particular interest in the context of future catalytic applications of AuAg alloy NCs. This work furthers understanding into the structure-property relationship and alloying behavior of large AuAg alloy NCs, which will be applied in future discovery of AuAg NCs for use in various applications.

#### **Author contributions**

David Morris: conceptualization, methodology, investigation, XAS data collection, XAS data analysis, visualization, data curation, formal analysis, validation, writing - original draft, writing - review and editing. Xiangsha Du: sample preparation, mass spectrometry data collection, mass spectrometry data analysis, writing - review and editing. Rongchao Jin: supervision, writing - review and editing, resources, funding acquisition. Peng Zhang: supervision, writing - review and editing, resources, funding acquisition.

## Data availability

The data supporting this article has been included as part of the ESI.†

### Conflicts of interest

There are no conflicts to declare.

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